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ENTRY SESSION 0.49

TOTAL

0.71

SESSION WILL BE HELD FOR 120 MINUTES STN INTERNATIONAL SESSION SUSPENDED AT 08:15:42 ON 25 JUN 2010

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 SINCE FILE
 TOTAL

 SENTRY
 SESSION

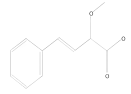
 FULL ESTIMATED COST
 0.49
 0.71

=>

Uploading C:\Documents and Settings\PZucker\My Documents\Examination Auxillary files\10566995\10566995 AF PHENYLBUTENOIC ACIDS.str

L1 STRUCTURE UPLOADED

=> D L1 L1 HAS NO ANSWERS L1 STR



Structure attributes must be viewed using STN Express query preparation.

=> SEARCH L1 SSS SAM
SAMPLE SEARCH INITIATED 08:24:11 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 323 TO ITERATE

100.0% PROCESSED 323 ITERATIONS SEARCH TIME: 00.00.01 1 ANSWERS

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**
PROJECTED ITERATIONS: 5382 TO 7538
PROJECTED ANSWERS: 1 TO 80

L2 1 SEA SSS SAM L1

=> D SCAN

L2 1 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
IN 3-Butenoic acid, 2-(acetyloxy)-4-phenyl-, (2R,3E)MF C12 H12 O4

Absolute stereochemistry. Rotation (-). Double bond geometry as shown.

ALL ANSWERS HAVE BEEN SCANNED

=> SEARCH L1 SSS FULL FULL SEARCH INITIATED 08:24:46 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 6297 TO ITERATE

100.0% PROCESSED 6297 ITERATIONS SEARCH TIME: 00.00.01 35 ANSWERS

L3 35 SEA SSS FUL L1

=> DSCAN

L4 0 DSCAN

=> D SCAN L3

L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN IN 3-Butenoic acid, 2-(acetyloxy)-4-phenyl-, (2S,3E)-MF C12 H12 04

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):10

L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
3-Butenoic acid, 4-phenyl-2-(phenylmethoxy)-,
(3R)-tetrahydro-4,4-dimethyl-2-oxo-3-furanyl ester, (2S,3E)MF C23 H24 05

Absolute stereochemistry. Double bond geometry as shown.

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic-2-d acid, 2-ethoxy-4-phenyl-, ethyl ester, (E)- (9CI)
- MF C14 H17 D O3

Double bond geometry as shown.

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic acid, 4-phenyl-2-[2-(trimethylsilyl)ethoxy]-, methyl ester, (2S)-
- MF C16 H24 O3 Si

Absolute stereochemistry. Double bond geometry unknown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN Benzeneacetic acid, α -methoxy- α -(trifluoromethyl)-,
- (2E)-1-(ethoxycarbonyl)-3-phenyl-2-propen-1-yl ester, (αR)-
- MF C22 H21 F3 O5

Absolute stereochemistry. Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic acid, 2-[[[bis(1-methylethyl)amino]carbonyl]oxy]-4-phenyl-, methyl ester, (2R,3E)-
- MF C18 H25 N O4

Absolute stereochemistry. Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic acid, 4-phenyl-2-(phenylmethoxy)-, methyl ester, (3E)-
- MF C18 H18 O3

Double bond geometry as shown.

- 1.3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN Benzeneacetic acid, α -methoxy- α -(trifluoromethyl)-,
- (1S, 2E)-1-(ethoxycarbonyl)-3-phenyl-2-propen-1-yl ester, (αR)-

C22 H21 F3 O5 MF

Absolute stereochemistry. Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic acid, 2-(2-chloro-1,3-dioxobutoxy)-4-phenyl-, methyl ester, (2R, 3E)-
- MF C15 H15 C1 O5

Absolute stereochemistry. Double bond geometry as shown.

- 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic acid, 2-[(3-methyl-2-buten-1-yl)oxy]-4-phenyl-, methyl ester, (3E) -
- MF C16 H20 O3

Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN IN 3-Butenoic acid, 2-methoxy-4-phenyl-, methyl ester C12 H14 O3 MF
- O OMe MeO-C-CH-CH-CH-Ph

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):25

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN TN
- 3-Butenoic acid, 2-methoxy-4-phenyl-, methyl ester, (3E)-C12 H14 O3 ME

Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- T. 3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- Benzeneacetic acid, α -methoxy- α -(trifluoromethyl)-, IN
- 1-(methoxycarbonyl)-3-phenyl-2-propenyl ester, [R-[R*,R*-(E)]]- (9CI) MF C21 H19 F3 O5

Absolute stereochemistry. Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN L3
- IN 3-Butenoic acid, 4-phenyl-2-propoxy-
- C13 H16 O3

ME

OPr-n

Ph-CH-CH-CH-CO2H

- **PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT**
- 1.3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- TN 3-Butenoic acid, 2-[(2E)-2-hexen-1-yloxy]-4-phenyl-, 1-methylethyl ester
- MF C19 H26 O3

Double bond geometry as described by E or Z.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

T. 3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN 3-Butenoic acid, 2-(acetyloxy)-4-phenyl-, (S)- (9CI) TN C12 H12 O4 MF

Absolute stereochemistry. Double bond geometry unknown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- 1.3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic acid, 2-(acetyloxy)-4-phenyl-, (3E)-
- C12 H12 O4 MF

Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic acid, 2-methoxy-4-phenyl-, (3S)-tetrahydro-4, 4-dimethyl-2-oxo-3-furanyl ester, (2R, 3E)-
- C17 H20 O5

Absolute stereochemistry. Double bond geometry as shown.

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic acid, 4-phenyl-2-(2,2,2-trifluoroethoxy)-, ethyl ester, (E)-(9CI)
- MF C14 H15 F3 O3

Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic acid, 2-(acetyloxy)-4-phenyl-, (2R,3E)-MF C12 H12 O4

Absolute stereochemistry. Rotation (-). Double bond geometry as shown.

- **PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT**
- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic acid, 4-phenyl-2-(2-propen-1-yloxy)-,
- (3R)-tetrahydro-4,4-dimethyl-2-oxo-3-furanyl ester, (2S,3E)-MF C19 H22 O5

... 020 00

Absolute stereochemistry. Double bond geometry as shown.

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic acid, 2-(acetyloxy)-4-phenyl-, methyl ester, (2S,3E)MF C13 H14 O4

Absolute stereochemistry. Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic acid, 2-methoxy-4-phenyl-, (3E)-
- MF C11 H12 O3

Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN Benzeneacetic acid, α-methoxy-α-(trifluoromethyl)-,
 - (1R, 2E)-1-(ethoxycarbonyl)-3-phenyl-2-propen-1-yl ester, (αR)-
- MF C22 H21 F3 O5

Absolute stereochemistry.

Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic acid, 2-(1,3-dioxobutoxy)-4-phenyl-, methyl ester, (2R,3E)-MF C15 H16 O5

Absolute stereochemistry. Rotation (-). Double bond geometry as shown.

- **PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT**
- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic acid, 2-(acetyloxy)-4-phenyl-, ethyl ester, (3E)-
- MF C14 H16 O4

Double bond geometry as shown.

- **PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT**
- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN

IN 3-Butenoic acid, 4-phenyl-2-(phenylmethoxy)-, methyl ester MF C18 H18 O3

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic acid, 2-[[2-chloro-2-[2-(4chlorophenyl)hydrazinylidene]acetyl]oxy]-4-phenyl-, methyl ester, (2R,3E)-MF C19 H16 C12 N2 O4

Absolute stereochemistry.
Double bond geometry as described by E or Z.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN Benzeneacetic acid, α-methoxy-α-(trifluoromethyl)-,
 - 1-(methoxycarbonyl)-3-phenyl-2-propenyl ester, [S-[R*,S*-(E)]]- (9CI)
- MF C21 H19 F3 O5

Absolute stereochemistry. Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic acid, 2-methoxy-4-phenyl-
- MF C11 H12 O3

ОМе

Ph-CH-CH-CH-CO2H

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic acid, 2-[(3-methyl-2-buten-1-yl)oxy]-4-phenyl-, 1-methylethyl ester
- MF C18 H24 O3

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic acid, 2-methoxy-4-phenyl-,
- tetrahydro-4,4-dimethyl-2-oxo-3-furanyl ester, $[R-[R^*,S^*-(E)]]$ (9CI) MF C17 H20 O5

Absolute stereochemistry. Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic acid, 4-phenyl-2-propoxy-, methyl ester

- L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN 3-Butenoic acid, 2-[(2-methyl-2-propen-1-yl)oxy]-4-phenyl-, 1-methylethyl ester

MF C17 H22 O3

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L3 35 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN IN 3-Butenoic acid, 2-ethoxy-4-phenyl-, ethyl ester, (E)- (9CI) MF C14 H18 03

Double bond geometry as shown.

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SINCE FILE TOTAL
ENTRY SESSION
FULL ESTIMATED COST 198.51 198.73

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REVISED CLASS FIELDS (/NCL) LAST RELOADED: Apr 2010
USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Apr 2010

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=> L3 L5 21 L3 => D L5 1-21 TI

- L5 ANSWER 1 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- TI Enantioselective C-C Bond Formation by Rhodium-Catalyzed Tandem Ylide Formation/[2,3]-Sigmatropic Rearrangement between Donor/Acceptor Carbenoids and Allylic Alcohols
- L5 ANSWER 2 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- $\hbox{ \it TI Pincer Complex-Catalyzed Redox Coupling of Alkenes with Iodonium Salts via Presumed Palladium(IV) Intermediates } \\$
- L5 ANSWER 3 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- TI Influence of electron-deficient ruthenium(I) carbonyl carboxylates on the vinylogous reactivity of metal carbenoids
- L5 ANSWER 4 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- TI Regioselectivity in Lewis acids catalyzed X-H (O, S, N) insertions of methyl styryldiazoacetate with benzyl alcohol, benzyl thiol, and aniline
- L5 ANSWER 5 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- TI Practical Approach to $\alpha-$ or $\gamma-$ Heterosubstituted Enoic Acids
- L5 ANSWER 6 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- TI Catalytic Enantioselective O-H Insertion Reactions
- L5 ANSWER 7 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- TI Chemo-enzymatic synthesis of (R)- and (S)-2-hydroxy-4-phenylbutanoic acid via enantio-complementary deracemization of
 - (±)-2-hydroxy-4-phenyl-3-butenoic acid using a racemase-lipase two-enzyme system

- L5 ANSWER 8 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- TI Preparation of 4-phenyl-but-3-enoic acid derivatives, as peroxisome proliferator-activated receptors (PPAR) ligands, in particular PPAR α and PPAR γ agonists, for the treatment and prevention of diabetes, dyslipidemia, atherosclerosis
- L5 ANSWER 9 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- TI Asymmetric reduction of alkyl 2-oxo-4-arylbutanoates and -but-3-enoates by Candida parapsilosis ATCC 7330: assignment of the absolute configuration of ethyl 2-hydroxy-4-(p-methylphenyl)but-3-enoate by 1H NMR
- L5 ANSWER 10 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- TI Three-Component Reaction of Aryl Diazoacetates, Alcohols, and Aldehydes (or Imines): Evidence of Alcoholic Oxonium Ylide Intermediates
- 5 ANSWER 11 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- TI Chiral catalyst enhancement of diastereocontrol for O-H insertion reactions of styryl- and phenyldiazoacetate esters of pantolactone
- L5 ANSWER 12 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- II Synthesis of α -allyloxy-substituted α, β -unsaturated esters via aldol condensation. Convenient access to highly substituted allyl vinyl ethers
- L5 ANSWER 13 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- TI Regiochemistry of molybdenum-catalyzed O-H insertions of vinylcarbenoids
- L5 ANSWER 14 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- TI The first case of asymmetric induction in intramolecular nitrile imine cycloadditions: synthesis of enantiopure 3-substituted 6-oxo-2,3,3a,5-tetrahydro-4-carbomethoxy-furo[3,4-c]pyrazoles
- L5 ANSWER 15 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- TI Enantioselective lithiation and substitution of (E)-cinnamyl N,N-diisopropylcarbamate through use of (-)-sparteine complexes
- L5 ANSWER 16 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- TI Chemoenzymic synthesis of enantiomerically pure alkene 1,2-diols and glycosides thereof
- L5 ANSWER 17 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- TI A Stereospecific Access to Allylic Systems Using Rhodium(II)-Vinyl Carbenoid Insertion into Si-H, O-H, and N-H Bonds
- L5 ANSWER 18 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- TI Electronic versus steric effects in 5-endo-trig-like electrophilic cyclizations
- L5 ANSWER 19 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- TI Enzymic resolution of 2-hydroxy-4-phenylbutanoic acid and 2-hydroxy-4-phenylbutenoic acid
- L5 ANSWER 20 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- TI Rhodium(II)-vinylcarbenoid insertion into the Si-H bond. A new stereospecific synthesis of allylsilanes
- L5 ANSWER 21 OF 21 CAPLUS COPYRIGHT 2010 ACS on STN
- TI Enantioselective reduction of β, χ -unsaturated α -keto acids using Bacillus stearothermophilus lactate dehydrogenase: a new route to functionalized allylic alcohols

=> FILE REG COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 9.69 208.42

FULL ESTIMATED COST

L6 RN

ED

841202-04-8 REGISTRY Entered STN: 03 Mar 2005

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=> E 3-Butenoic acid, 2-methoxy-4-phenyl-/CN
E1
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             1
                  LO(3.3.1.13,7) DEC-2-YLAMINO) CARBONYL) -1H-PYRAZOL-1-YL) -/CN
E 2
                  3-BUTENOIC ACID, 2-METHOXY-4-(TRIMETHYLSILYL)-3-((TRIMETHYLS
                  ILYL)OXY)-, TRIMETHYLSILYL ESTER, (E)-/CN
             1 --> 3-BUTENOIC ACID, 2-METHOXY-4-PHENYL-/CN
E4
                  3-BUTENOIC ACID, 2-METHOXY-4-PHENYL-, (3E)-/CN
E5
                  3-BUTENOIC ACID, 2-METHOXY-4-PHENYL-, (3S)-TETRAHYDRO-4,4-DI
             1
                  METHYL-2-OXO-3-FURANYL ESTER, (2R,3E)-/CN
                 3-BUTENOIC ACID, 2-METHOXY-4-PHENYL-, METHYL ESTER/CN
E6
            1
E7
            1
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E8
                  3-BUTENOIC ACID, 2-METHOXY-4-PHENYL-, TETRAHYDRO-4,4-DIMETHY
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                  3-BUTENOIC ACID, 2-METHOXYETHYL ESTER/CN
E9
E10
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            1
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E11
            1
                  YL)METHYL ESTER/CN
E12
            1
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                  TER, S,S-DIOXIDE/CN
=> E3
L6
            1 "3-BUTENOIC ACID, 2-METHOXY-4-PHENYL-"/CN
=> D L6
    ANSWER 1 OF 1 REGISTRY COPYRIGHT 2010 ACS on STN
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3-Butenoic acid, 2-methoxy-4-phenyl- (CA INDEX NAME)

OTHER NAMES: CN 2-Methoxy-4-pheny1but-3-enoic acid MF C11 H12 O3 SR CA

LC STN Files: CA, CAPLUS, CASREACT, USPAT2, USPATFULL

OMe

Ph-CH-CH-CO2H

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1 REFERENCES IN FILE CA (1907 TO DATE)
1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

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=> L6 L7 1 L6

- L7 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2010 ACS on STN
- AN 2005:119915 CAPLUS <<LOGINID::20100625>>
- DN 142:219047
- TI Preparation of 4-phenyl-but-3-enoic acid derivatives, as peroxisome proliferator-activated receptors (PPAR) ligands, in particular PPAR α and PPAR γ agonists, for the treatment and prevention of diabetes, dyslipidemia, atherosclerosis
- IN Zeiller, Jean Jacques; Dumas, Herve; Guyard Dangremont, Valerie; Berard, Isabelle; Contard, Francis; Guerrier, Daniel; Ferrand, Gerard; Bonhomme, Yves
- PA Merck Sante, Fr. SO Fr. Demande, 38 p
 - Fr. Demande, 38 pp. CODEN: FRXXBL
- DT Patent
- LA French

| FAN. | CNT | 1 | | | | | | | | | | | | | | | | | |
|------|------------|--------------|------------|------|------|-------|---|----------|--------|-----|--|------|------|-------|----------|----------|------|-----|--|
| | PATENT NO. | | | | | | | DATE | | | APPLICATION NO. | | | | | | | | |
| PI | | 2858 | | | | | | | | | FR 2 | | | | | | 0030 | 804 | |
| | FR | 2858 | 615 | | | B1 | | 2006 | 1222 | | | 000 | | | | _ | | | |
| | AII | 2004 | 2632 | 54 | | A1 | | 2005 | 0217 | | AII 2 | 004- | 2632 | 5.4 | | 2 | 0040 | 714 | |
| | AU | 2004 | 2004263254 | | | B2 | | 20100318 | | | | | | | | | | | |
| | CA | 2534 | 2534493 | | | A1 | | 20050217 | | | CA 2004-2534493 | | | | | 20040714 | | | |
| | | | | | | | | | | | WO 2004-EP7776 | | | | | | | | |
| | | W: | AE, | AG, | AL, | AM, | AT, | AU, | AZ. | BA, | BB, | BG, | BR, | BW, | BY, | BZ, | CA, | CH, | |
| | | | CN, | co, | CR, | CU, | CZ, | DE, | DK, | DM, | DZ, | EC, | EE, | EG, | ES, | FI, | GB, | GD, | |
| | | | GE, | GH, | GM, | HR, | HU, | ID, | IL, | IN, | IS, | JP, | KE, | KG, | KP, | KR, | KZ, | LC, | |
| | | | LK, | LR, | LS, | LT, | LU, | LV, | MA, | MD, | MG, | MK, | MN, | MW, | MX, | MZ, | NA, | NI, | |
| | | | NO, | NZ, | OM, | PG, | PH, | PL, | PT, | RO, | RU, | SC, | SD, | SE, | SG, | SK, | SL, | SY, | |
| | | | TJ, | TM, | TN, | TR, | TT, | TZ, | UA, | UG, | US, | UZ, | VC, | VN, | YU, | ZA, | ZM, | zw | |
| | | RW: | BW, | GH, | GM, | KE, | LS, | MW, | MZ, | NA, | SD, | SL, | SZ, | TZ, | UG, | ZM, | ZW, | AM, | |
| | | | AZ, | BY, | KG, | KZ, | MD, | RU, | ТJ, | TM, | AT, | BE, | BG, | CH, | CY, | CZ, | DE, | DK, | |
| | | | | | | | | | | | IT, | | | | | | | | |
| | | | | | | | ВJ, | CF, | CG, | CI, | CM, | GΑ, | GN, | GQ, | GW, | ML, | MR, | NE, | |
| | | | | TD, | | | | | | | | | | | | | | | |
| | EP | | | | | | | | | | EP 2004-740992
B, GR, IT, LI, LU, NL, | | | | | | | | |
| | | R: | | | | | | | | | | | | | | | MC, | PT, | |
| | | | | | | | | | | | BG, | | | | | | | | |
| | JP | P 2007501190 | | | | Т | T 20070125 JP 2006-522255
A1 20060810 US 2006-566995 | | | | | | | 55 | 20040714 | | | | |
| | | | | | | | | | | | US 2 | 006- | 5669 | 95 | | 2 | 0060 | 202 | |
| PRAI | | 2003 | | | | | | | | | | | | | | | | | |
| | | 2004 | | | | | | | | | | | | | | _ | | | |
| | | ENT H | | | | | | | | | N LS | US D | ISPL | AY F | ORMA | T | | | |
| | | SREAC' | | | | | | | | | 2212 T | D. | E E0 | D TII | TC D | ECOD | D | | |
| RE.C | IN I | 12 | | | | | | | | | RE | | | K IH | 19 K | LCOK | ע | | |
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=> FIE REG 174 FIE 14 FIES 188 FIE

(FIE OR FIES) 1734 REG 139 REGS

1846 REG

(REG OR REGS)
L8 0 FIE REG
(FIE(W)REG)

REGISTRY INITIATED

Substance data EXPAND from CAS REGISTRY in progress...

| E1 | 1 | 3-BUTENOIC ACID, 4-PHENYL-2-((TRIPHENYLMETHYL)AMINO)-, (E)-/CN |
|-----|----|---|
| E2 | 1 | 3-BUTENOIC ACID, 4-PHENYL-2-(1-PHTHALAZINYLHYDRAZONO)-/CN |
| E3 | 1> | 3-BUTENOIC ACID, 4-PHENYL-2-(2,2,2-TRIFLUOROETHOXY)-, ETHYL ESTER, (E)-/CN |
| E 4 | 1 | 3-BUTENOIC ACID, 4-PHENYL-2-(2,2,2-TRIMETHYL-1,1-BIS(TRIMETH YLSILYL)DISILANYL)-, ETHYL ESTER, (E)-/CN |
| E5 | 1 | 3-BUTEMOIC ACID, 4-PHENYL-2-(2,2,2-TRIMETHYL-1,1-BIS(TRIMETH YLSILYL)DISIANYL)-, TETRAHYDRO-4,4-DIMETHYL-2-OXO-3-FURANYL ESTER, (5-(R*,5*-(E)))-/CN |
| E6 | 1 | 3-BUTENOIC ACID, 4-PHENYL-2-(2-((PHENYLAMINO)CARBONYL)HYDRAZ INYLIDENE)-/CN |
| E7 | 1 | 3-BUTENOIC ACID, 4-PHENYL-2-(2-((PHENYLAMINO)CARBONYL)HYDRAZ INYLIDENE)-, HYDRAZIDE/CN |
| E8 | 1 | 3-BUTENOIC ACID, 4-PHENYL-2-(2-((PHENYLAMINO)THIOXOMETHYL)HY
DRAZINYLIDENE)-/CN |
| E9 | 1 | 3-BUTENOIC ACID, 4-PHENYL-2-(2-(1-PHTHALAZINYL)HYDRAZINYLIDE NE)-/CN |
| E10 | 1 | 3-BUTENOIC ACID, 4-PHENYL-2-(2-(4-QUINAZOLINYL)HYDRAZINYLIDE
NE)-/CN |
| E11 | 1 | 3-BUTENOIC ACID, 4-PHENYL-2-(2-(TRIFLUOROMETHYL)PHENOXY)-, (3E)-/CN |
| E12 | 1 | 3-BUTENOIC ACID, 4-PHENYL-2-(2-(TRIFLUOROMETHYL)PHENOXY)-, METHYL ESTER, (3E)-/CN |

=> E3

REG1stRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

```
L10 2 L9
```

=> D L10 1-2 TI

- L10 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2010 ACS on STN
- TI A Stereospecific Access to Allylic Systems Using Rhodium(II)-Vinyl Carbenoid Insertion into Si-H, O-H, and N-H Bonds
- L10 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2010 ACS on STN
- TI Electronic versus steric effects in 5-endo-trig-like electrophilic cyclizations

=> D L10 FILE REG

'FILE' IS NOT A VALID FORMAT FOR FILE 'CAPLUS' 'REG' IS NOT A VALID FORMAT FOR FILE 'CAPLUS'

The following are valid formats:

```
ABS ----- GI and AB
ALL ----- BIB, AB, IND, RE
APPS ----- AI, PRAI
BIB ----- AN, plus Bibliographic Data and PI table (default)
CAN ----- List of CA abstract numbers without answer numbers
CBIB ----- AN, plus Compressed Bibliographic Data
CLASS ----- IPC, NCL, ECLA, FTERM
DALL ---- ALL, delimited (end of each field identified)
DMAX ----- MAX, delimited for post-processing
FAM ----- AN, PI and PRAI in table, plus Patent Family data
FBIB ----- AN, BIB, plus Patent FAM
IND ----- Indexing data
IPC ----- International Patent Classifications
MAX ----- ALL, plus Patent FAM, RE
PATS ----- PI, SO
SAM ----- CC, SX, TI, ST, IT
SCAN ----- CC, SX, TI, ST, IT (random display, no answer numbers;
             SCAN must be entered on the same line as the DISPLAY,
             e.g., D SCAN or DISPLAY SCAN)
STD ---- BIB, CLASS
IABS ----- ABS, indented with text labels
IALL ----- ALL, indented with text labels
IBIB ----- BIB, indented with text labels
IMAX ----- MAX, indented with text labels
ISTD ----- STD, indented with text labels
OBIB ----- AN, plus Bibliographic Data (original)
OIBIB ----- OBIB, indented with text labels
SBIB ----- BIB, no citations
SIBIB ----- IBIB, no citations
HIT ----- Fields containing hit terms
HITIND ----- IC, ICA, ICI, NCL, CC and index field (ST and IT)
             containing hit terms
HITRN ----- HIT RN and its text modification
HITSTR ----- HIT RN, its text modification, its CA index name, and
             its structure diagram
HITSEO ----- HIT RN, its text modification, its CA index name, its
             structure diagram, plus NTE and SEO fields
FHITSTR ---- First HIT RN, its text modification, its CA index name, and
             its structure diagram
FHITSEQ ---- First HIT RN, its text modification, its CA index name, its
             structure diagram, plus NTE and SEQ fields
KWIC ----- Hit term plus 20 words on either side
OCC ----- Number of occurrence of hit term and field in which it occurs
```

To display a particular field or fields, enter the display field codes. For a list of the display field codes, enter HELD DFIELDS at an arrow prompt (=>). Examples of formats include: TI, TI, AU, BIB, ST, TI, IND; TI, SO. You may specify the format fields in any order and the information will be displayed in the same order as the format specification.

All of the formats (except for SAM, SCAN, HIT, HITIND, HITRN, HITSTR, FHITSTR, HITSEQ, FHITSEQ, KWIC, and OCC) may be used with DISPLAY ACC to view a specified Accession Number.

ENTER DISPLAY FORMAT (BIB): END

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 2.28 233.69

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STRUCTURE FILE UPDATES: 24 JUN 2010 HIGHEST RN 1228341-10-3 DICTIONARY FILE UPDATES: 24 JUN 2010 HIGHEST RN 1228341-10-3

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http://www.cas.org/support/stngen/stndoc/properties.html

FS

ME

STEREOSEARCH

C14 H18 O3 SR CA

```
=> E 3-Butenoic acid, 2-ethoxy-4-phenyl-, ethyl ester, (E)-/CN
E1
            1
                 3-BUTENOIC ACID, 2-ETHOXY-4, 4-DIPHENYL-, ETHYL ESTER/CN
E2
                  3-BUTENOIC ACID, 2-ETHOXY-4-(4-(PHENYLMETHOXY)PHENYL)-, ETHY
            1
                  L ESTER/CN
EЗ
            1 --> 3-BUTENOIC ACID, 2-ETHOXY-4-PHENYL-, ETHYL ESTER, (E)-/CN
            1 3-BUTENOIC ACID, 2-ETHOXYETHYL ESTER/CN
1 3-BUTENOIC ACID, 2-ETHYL-/CN
E4
E5
E6
            1
                 3-BUTENOIC ACID, 2-ETHYL-, (1S,3S)-3-((4-METHOXYPHENYL)METHO
                 XY)-1-(1-METHYLETHENYL)BUTYL ESTER, (2R)-/CN
E7
            1
                 3-BUTENOIC ACID, 2-ETHYL-, (1S,3S)-3-((4-METHOXYPHENYL)METHO
                 XY)-1-(1-METHYLETHENYL)BUTYL ESTER, (2S)-/CN
                 3-BUTENOIC ACID, 2-ETHYL-, (2R)-/CN
E8
E9
           1
                 3-BUTENOIC ACID, 2-ETHYL-, 1,1-DIMETHYLETHYL ESTER, (2R)-/CN
E10
           1
                 3-BUTENOIC ACID, 2-ETHYL-, 1-METHYLETHYL ESTER, (2R)-/CN
E11
           2
                 3-BUTENOIC ACID, 2-ETHYL-, 2'-HYDROXY(1,1'-BINAPHTHALEN)-2-Y
                 L ESTER, STEREOISOMER/CN
E12
           1
                 3-BUTENOIC ACID, 2-ETHYL-, 2-(4-BROMOPHENYL)-2-OXOETHYL ESTE
                  R/CN
=> E3
            1 "3-BUTENOIC ACID, 2-ETHOXY-4-PHENYL-, ETHYL ESTER, (E)-"/CN
=> D L11
L11 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2010 ACS on STN
    173602-52-3 REGISTRY
RN
ED
    Entered STN: 28 Feb 1996
CN 3-Butenoic acid, 2-ethoxy-4-phenyl-, ethyl ester, (E)- (9CI)
    (CA INDEX NAME)
```

Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1907 TO DATE)
2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

=> FILE CAPLUS COST IN U.S. DOLLARS FULL ESTIMATED COST

SINCE FILE TOTAL ENTRY SESSION 8.09 241.78

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FILE COVERS 1907 - 25 Jun 2010 VOL 153 ISS 1 FILE LAST UPDATED: 24 Jun 2010 (20100624/ED) REVISED CLASS FIELDS (/NCL) LAST RELOADED: Apr 2010 USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Apr 2010

CAplus now includes complete International Patent Classification (IPC) reclassification data for the second quarter of 2010.

CAS Information Use Policies apply and are available at:

http://www.cas.org/legal/infopolicy.html

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> L11 L12 2 L11

=> D L12 1-2 TI FBIB ABS

- L12 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2010 ACS on STN
- A Stereospecific Access to Allylic Systems Using Rhodium(II)-Vinyl Carbenoid Insertion into Si-H, O-H, and N-H Bonds

AN 1997:198048 CAPLUS <<LOGINID::20100625>>

- 126:211638 DM
- OREF 126:40925a,40926a
- A Stereospecific Access to Allylic Systems Using Rhodium(II)-Vinyl
- Carbenoid Insertion into Si-H, O-H, and N-H Bonds AU Bulugahapitiva, Privadarshanie; Landais, Yannick; Parra-Rapado, Liliana;
- Planchenault, Denis; Weber, Valery CS College Propedeutique, Universite de Lausanne, Lausanne-Dorigny, 1015,
- Switz. SO Journal of Organic Chemistry (1997), 62(6), 1630-1641
- CODEN: JOCEAH; ISSN: 0022-3263 PB American Chemical Society
- DT Journal
- LA English
- AB Rhodium-catalyzed decomposition of a-vinyl diazo esters in the presence of silanes, alcs., ethers, amines, and thiols has been shown to produce the corresponding a-silyl, a-hydroxy, a-alkoxy,

α-amino, and α-thioalkoxy esters in generally good yield with

a complete retention of the stereochem, of the double bond of the diazo precursor. An extension of the process in homochiral series has also been devised using either a chiral auxiliary attached to the ester function or achiral α-vinyl diazo esters and Doyle's chiral catalyst Rh2(MEPY) 4.

In the former approach, pantolactone as chiral auxiliary gave diastereoselectivities of up to 70%, while the second approach produced the desired allylsilane with ee as high as 72%. On the other hand, Rh2(MEPY)4-catalyzed insertion into the O-H bond of water led to poor or

no enantioselectivity in good agreement with recent literature reports. OSC.G 59 THERE ARE 59 CAPLUS RECORDS THAT CITE THIS RECORD (60 CITINGS) RE.CNT 119 THERE ARE 119 CITED REFERENCES AVAILABLE FOR THIS RECORD

- L12 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2010 ACS on STN
- ΤI Electronic versus steric effects in 5-endo-trig-like electrophilic cyclizations

ALL CITATIONS AVAILABLE IN THE RE FORMAT

- AN 1995:974892 CAPLUS <<LOGINID::20100625>>
- DN 124:176328 OREF 124:32707a,32710a
- TI Electronic versus steric effects in 5-endo-trig-like electrophilic cvclizations
- ΑU Landais, Yannick; Planchenault, Denis
- CS Inst. de Chimie Organique, Univ. de Lausanne, Lausanne-Dorigny, 1015,
- Synlett (1995), (11), 1191-3 SO CODEN: SYNLES; ISSN: 0936-5214
- Switz. Thieme PB
- DT Journal LA English
- CASREACT 124:176328 OS



AB Electronically and sterically differentiated allylic substituents such as RO, NHPh, PhS, and PhSO2 groups were used to demonstrate the influence of electronic and/or steric effects in the stereocontrol of the PhSeC1-promoted electrophilic 3-endo-trig-like cyclizations of 2-substituted-3-alkenols, (E)-PhCH:CHCHXCH20H (1, X = OH, OEH, OCH2CF3, OPh, NHPh, SPh). 1 Reacted with PhSeC1/K2CO3 to give predominantly the 2,4-trans-tetrahydrofuran I, however, the cis-2,4-diastereoisomer II was predominant for X = NHPh and SPh for reasons of electronic effects.

OSC.G 20 THRE ARE 20 CAPLUS RECORDS THAT CITE THIS RECORD (20 CITINGS)

=> FILE REG COST IN U.S. DOLLARS SINCE FILE TOTAL. ENTRY SESSION FULL ESTIMATED COST 8.20 249.98 SINCE FILE DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) TOTAL ENTRY SESSION -1.70 -1.70 CA SUBSCRIBER PRICE

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TSCA INFORMATION NOW CURRENT THROUGH January 8, 2010.

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http://www.cas.org/support/stngen/stndoc/properties.html

```
=> e 3-Butenoic acid, 2-(acetyloxy)-4-phenyl-, ethyl ester,/cn
                  3-BUTENOIC ACID, 2-(ACETYLOXY)-4-PHENYL-, (3E)-/CN
E1
E2
                  3-BUTENOIC ACID, 2-(ACETYLOXY)-4-PHENYL-, (S)-/CN
             0 --> 3-BUTENOIC ACID, 2-(ACETYLOXY)-4-PHENYL-, ETHYL ESTER,/CN
E3
Ε4
             1
                  3-BUTENOIC ACID, 2-(ACETYLOXY)-4-PHENYL-, ETHYL ESTER, (3E)-
E5
                  3-BUTENOIC ACID, 2-(ACETYLOXY)-4-PHENYL-, METHYL ESTER, (2S,
                  3E) -/CN
                  3-BUTENOIC ACID, 2-(ACETYLOXY)-4-PHENYL-, METHYL ESTER, (S-(
                  E))-/CN
E7
                  3-BUTENOIC ACID, 2-(ACETYLOXY)ETHYL ESTER/CN
E8
            1
                  3-BUTENOIC ACID, 2-(ACETYLTHIO)-2-METHYL-, PHENYLMETHYL ESTE
                  R, (2R)-/CN
E9
           1
                 3-BUTENOIC ACID, 2-(ACETYLTHIO)-2-METHYL-, PHENYLMETHYL ESTE
```

```
R, (2S)-/CN
E10
                   3-BUTENOIC ACID, 2-(ACETYLTHIO)-2-METHYL-4-PHENYL-, PHENYLME
                   THYL ESTER, (2R, 3E) -/CN
E11
             1
                   3-BUTENOIC ACID, 2-(ACETYLTHIO)-2-METHYL-4-PHENYL-, PHENYLME
                   THYL ESTER, (2S, 3E) -/CN
                   3-BUTENOIC ACID, 2-(AMINO((2-AMINOPHENYL)AMINO)METHYLENE)-4,
E12
             1
                   4-DICYANO-, ETHYL ESTER, (Z)-/CN
=> e4
L13
             1 "3-BUTENOIC ACID, 2-(ACETYLOXY)-4-PHENYL-, ETHYL ESTER, (3E)-"/C
=> d 113
L13 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2010 ACS on STN
     1151782-13-6 REGISTRY
RN
ED
     Entered STN: 02 Jun 2009
CN
     3-Butenoic acid, 2-(acetyloxy)-4-phenyl-, ethyl ester, (3E)-
     (CA INDEX NAME)
FS
     STEREOSEARCH
     C14 H16 O4
MF
SR
     CA
LC
     STN Files: CA, CAPLUS, CASREACT
Double bond geometry as shown.
          OAc
                OEt
Ph
**PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT**
               1 REFERENCES IN FILE CA (1907 TO DATE)
               1 REFERENCES IN FILE CAPLUS (1907 TO DATE)
=> e5
L14
             1 "3-BUTENOIC ACID, 2-(ACETYLOXY)-4-PHENYL-, METHYL ESTER, (2S,3E)
               - " / CN
=> d 114
L14 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2010 ACS on STN
RN
     206257-88-7 REGISTRY
     Entered STN: 03 Jun 1998
ED
     3-Butenoic acid, 2-(acetyloxy)-4-phenyl-, methyl ester, (2S,3E)-
     (CA INDEX NAME)
OTHER CA INDEX NAMES:
CN
     3-Butenoic acid, 2-(acetyloxy)-4-phenyl-, methyl ester, [S-(E)]-
FS
     STEREOSEARCH
```

Absolute stereochemistry.

STN Files: CA, CAPLUS

C13 H14 O4

MF

SR CA

Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1907 TO DATE)
1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

=> e6 L15 1 "3-BUTENOIC ACID, 2-(ACETYLOXY)-4-PHENYL-, METHYL ESTER, (S-(E)) -"/CN

=> d 115

- L15 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2010 ACS on STN
- RN 206257-88-7 REGISTRY
- ED Entered STN: 03 Jun 1998
- CN 3-Butenoic acid, 2-(acetyloxy)-4-phenyl-, methyl ester, (2S,3E)- (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 3-Butenoic acid, 2-(acetyloxy)-4-phenyl-, methyl ester, [S-(E)]-

- FS STEREOSEARCH
- MF C13 H14 O4
- SR CA
- LC STN Files: CA, CAPLUS

Absolute stereochemistry.

Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1907 TO DATE)
1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

=> file caplus
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SINCE FILE TOTAL
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ENTRY SESSION 0.00 -1.70

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FILE LAST UPDATED: 24 Jun 2010 (20100624/ED)
REVISED CLASS FIELDS (/NCL) LAST RELOADED: Apr 2010
USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Apr 2010

CAplus now includes complete International Patent Classification (IPC) reclassification data for the second quarter of 2010.

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> 113

L16 1 L13

=> d 116

L16 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2010 ACS on STN

AN 2009:652491 CAPLUS <<LOGINID::20100625>>

DN 151:8000

TI Pincer Complex-Catalyzed Redox Coupling of Alkenes with Iodonium Salts via Presumed Palladium(IV) Intermediates

AU Aydin, Juhanes; Larsson, Johanna M.; Selander, Nicklas; Szabo, Kalman J.
CS Department of Organic Chemistry, Arrhenius Laboratory, Stockholm

S Department of Organic Chemistry, Arrhenius Laboratory, Stockholm University, Swed.

SO Organic Letters (2009), 11(13), 2852-2854

CODEN: ORLEF7; ISSN: 1523-7060

PB American Chemical Society

DT Journal

LA English

OS CAŚREACT 151:8000

OSC.G 7 THERE ARE 7 CAPLUS RECORDS THAT CITE THIS RECORD (7 CITINGS)
RE.CNT 56 THERE ARE 56 CITED REFERENCES AVAILABLE FOR THIS RECORD

ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> 114

17 1 L14

```
=> d 117
L17 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2010 ACS on STN
AN 1998:217991 CAPLUS <<LOGINID::20100625>>
DN
     128:294949
OREF 128:58463a.58466a
    Chemoenzymic synthesis of enantiomerically pure alkene 1,2-diols and
     glycosides thereof
AU
     Ziegler, Thomas; Bien, Frank; Jurisch, Claus
CS
    Institute of Organic Chemistry, University of Cologne, Cologne, D-50939,
     Germany
SO
    Tetrahedron: Asymmetry (1998), 9(5), 765-780
     CODEN: TASYE3; ISSN: 0957-4166
PB
   Elsevier Science Ltd.
DT
    Journal
T.A
   English
os
   CASREACT 128:294949
OSC.G 29
             THERE ARE 29 CAPLUS RECORDS THAT CITE THIS RECORD (29 CITINGS)
RE.CNT 40
              THERE ARE 40 CITED REFERENCES AVAILABLE FOR THIS RECORD
              ALL CITATIONS AVAILABLE IN THE RE FORMAT
=> d 118
L18 NOT FOUND
The L-number entered has not been defined in this session, or it
has been deleted. To see the L-numbers currently defined in this
session, enter DISPLAY HISTORY at an arrow prompt (=>).
=> 5 115
MISSING OPERATOR 5 L15
The search profile that was entered contains terms or
nested terms that are not separated by a logical operator.
=> d 115
YOU HAVE REQUESTED DATA FROM FILE 'REGISTRY' - CONTINUE? (Y) /N:n
=> 115
L18
             1 L15
=> d 118
L18 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2010 ACS on STN
    1998:217991 CAPLUS <<LOGINID::20100625>>
AN
DN
     128:294949
OREF 128:58463a,58466a
ΤТ
     Chemoenzymic synthesis of enantiomerically pure alkene 1.2-diols and
     alvcosides thereof
ΔII
     Ziegler, Thomas: Bien, Frank: Jurisch, Claus
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     Institute of Organic Chemistry, University of Cologne, Cologne, D-50939,
SO
     Tetrahedron: Asymmetry (1998), 9(5), 765-780
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OSC.G 29

RE.CNT 40

CASREACT 128:294949

=> logoff hold
COST IN U.S. DOLLARS

FULL ESTIMATED COST

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)
CA SUBSCRIBER PRICE

SINCE FILE
ENTRY
ENTRY
SESSION
0.00
-1.70
-1.70

SESSION WILL BE HELD FOR 120 MINUTES
STN INTERNATIONAL SESSION SUSPENDED AT 08:50:46 ON 25 JUN 2010